MRI of the penis

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ABSTRACT. MRI of the penis is an expensive test that is not always superior to clinical examination or ultrasound. However, it shows many of the important structures, and in particular the combination of tumescence from intracavernosal alprostadil, and high-resolution $T_2$ sequences show the glans, corpora and the tunica albuginea well. In this paper we summarise the radiological anatomy and discuss the indications for MRI. For penile cancer, it may be useful in cases where the local stage is not apparent clinically. In priapism, it is an emerging technique for assessing corporal viability, and in fracture it can in most cases make the diagnosis and locate the injury. In some cases of penile fibrosis and Peyronie’s disease, it may aid surgical planning, and in complex pelvic fracture may replace or augment conventional urethrography. It is an excellent investigation for the malfunctioning penile prosthesis.

The penis is predominantly a superficial organ, and for the most part easily examined by palpation or with ultrasound. Although MRI can be used to assess most penile pathologies, it needs to demonstrate clinical utility to justify its expense. Before discussing the benefits of MRI for a number of pathologies, we will briefly summarise the radiological anatomy and scanning techniques.

Anatomy

The corpus spongiosum and corpora cavernosa are of high signal on $T_2$ weighted sequences, and intermediate to low on $T_1$ weighted sequences [1]. The tunica albuginea is a fibrous sheath surrounding all three (Figure 1), and is low in signal on all sequences, with the contrast between it and the higher signal corpora greatest on $T_2$ (Figures 2 and 3). The septum divides the two corpora cavernosa and is generally porous, allowing the passage of blood (and agents injected intracavernosally) from one side to the other [2].

Close to the tunica albuginea is Buck’s fascia, a slightly less dense fibrous sheath that is only sometimes distinguishable from the deeper tunica [3]. It fuses with the deep perineal fascia, so that haemorrhage from a penile fracture which does not compromise Buck’s fascia is confined to the penis, rather than spreading to the perineum. Superficial to Buck’s fascia is the thin, incomplete Dartos fascia (not clearly seen on MRI), then loose subepithelial connective tissue and skin. In the glans, the tunica albuginea is hard to distinguish from the more superficial subepithelial connective tissue and the two fuse towards the tip [4].

The deep parts of the corpora cavernosa are closely related to the inferior pubic rami on each side, and surrounded by the ischiocavernous muscle. The most proximal part of the corpus spongiosum is the bulb, which is surrounded by the bulbospongious muscle and is pierced by the urethra. The corpus spongiosum is continuous with the glans, and the two generally have the same signal characteristics on MRI.

The vascular anatomy of the penis is very variable [5], but the common penile artery, a branch of the internal pudendal artery, gives three relatively constant branches: crural, dorsal penile (supplying more distal corpus spongiosum and glans) and cavernosal, supplying the corpus cavernosum on each side and often duplicated. Side branches (the helicine arteries) are easily seen on ultrasound but are not usually visible on MRI. The venous anatomy is variable but the largest branches are the superficial and deep dorsal, in the midline, separated by Buck’s fascia.

MR sequence

The mainstay of penile MRI is high-resolution (matrix size $\approx$200), small-field-of-view, thin-slice (ideally 3 mm) $T_2$ weighted spin echo sequences in 3 orthogonal planes, with the penis in the anatomical position (resting on the anterior abdominal wall in the midline, although others place it dependently to minimise breathing artefact [6]). This is our routine protocol, with further sequences added as required. $T_1$ weighted images may be useful for the detection of haemorrhage and thrombosis within the corpora or draining veins. Small field-of-view short-tau inversion–recovery (STIR) sequences may show inflammation, and sometimes the deep veins of the pelvis well. For imaging the cavernosal vessels and corporal enhancement, we use dynamic contrast-enhanced sequences, with 3–4-mm slice thickness and a small field of view, usually in the coronal plane, as it shows the vessels in the base of the penis best. For assessment of
cavernosal viability in priapism, the dynamic scans show early perfusion, but there should be one scan at least 10 min after contrast—ideally a small field of view spin echo $T_1$ weighted sequence, with an identical sequence obtained before contrast for comparison. Contrast enhancement for staging tumours is usually not helpful [7].

**Tumescence and intracorporal agents**

There are two reasons for using intracavernosal agents to produce tumescence. First, the normal corpora become uniformly high in $T_2$ signal, and the tunica albuginea is seen as a thin, regular low signal layer around them. This makes it much easier to see both intracavernosal fibrosis and tunical plaque, as well as other intracavernosal pathologies such as fistula or tumour. Second, positioning of the tumescent (though ideally not fully so) penis in the midline makes it considerably easier to scan in true sagittal, coronal and axial planes. For young males with normal erectile function we use 5 mg of alprostadil, and for older males with erectile dysfunction the full dose of 20 mg. Priapism is rare, but can be treated in most cases by aspiration [8]. Sildenafil and manual stimulation may produce tumescence in most patients but, especially for MRI, they are not as reliable as intracavernosal agents [9].

**MRI for penile cancer**

The great majority of penile lesions are primary squamous cell tumours, although melanoma, basal cell carcinoma, sarcoma and lymphoma have been reported, and metastases are common enough to account for several cases series, with bladder the commonest site of origin—either haematogenous or from urethral spread of transitional cell carcinoma [10, 11]. Anterior urethral tumours are rare, and usually of the squamous rather than the transitional type [12].

Local staging (the classification is shown in Table 1) of penile cancer may be useful to plan the surgical approach. There is some controversy about the current classification, especially within the glans, which is the site of the majority of tumours [13] (Figure 4). It was noted in an early ultrasound study that the tunica albuginea in the glans becomes difficult to see and blends with subepithelial connective tissue [4]. Not only is the distinction between T1 and T2 disease therefore more difficult, but the finding of T2 disease in the glans has different implications for the same stage in the shaft.

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**Figure 1.** Cross-sectional penile anatomy.

**Figure 2.** $T_2$ (a) and $T_1$ (b) weighted sequences through the tumescent penis. Black arrowheads mark the tunica albuginea, and white arrowheads Buck’s fascia. In (a) the thick white arrow shows the superficial dorsal vein and the thinner white arrows the deep dorsal vessels. The cavernosal arteries are marked by black arrows. The urethra, lying in the middle of the corpus spongiosum, is marked by an asterisk.
with glansectomy (partial or complete) the treatment of choice [14, 15] as opposed to partial or complete penectomy and a significantly worse prognosis) in T2 disease involving the corpora cavernosa in the shaft [16]. The T3 classification is also problematic: involvement of the urethra is commonly near the meatus and may have a better prognosis than T2 involvement of the corpus spongiosum in the shaft [17]. These issues have led to the proposal of a new staging system more closely related to prognosis, also shown in Table 1 [18].

Tumour is seen as low signal compared with corpus spongiosum or cavernosum on $T_2$ sequences, but usually higher than the tunica (Figure 4). Although MRI provides excellent soft-tissue definition, it does not necessarily follow that it is superior to clinical examination for local staging, and early studies, usually without intracavernosal agents and in small numbers of patients, did not show definitively that it was of benefit. It must be borne in mind that the clinical stage is usually most accurately determined at operation and with the use of frozen section analysis. Later studies, with the use of intracavernosal agents, have shown that MRI was useful when infiltration of the corpora “could not be determined properly by clinical examination” and that MRI is probably better than ultrasound in this context [19]. In particular, several studies have shown that invasion of the corpora cavernosa is rarely missed [7, 19, 20], and proximal tumours, which are hard to scan on ultrasound and difficult to palpate, are shown well. However, even with intracorporal agents MRI can sometimes overstage tumours, and in one study of 55 patients 6 patients with T1 lesions were staged as T2 [20]. This was ascribed to technical factors—poor response to prostaglandin, previous radiotherapy, motion artefact and infection—but it is likely that some of the error was also due to fundamental limits to the resolution of MRI and the difficulty in distinguishing abutment and bulge from true invasion. No cases of priapism were seen, although in a previous smaller study priapism occurred in 1 out of 10 patients [7]. The use of contrast has been anecdotally described as useful [21], but the larger studies do not support its routine use [7]. For superficial lesions clinical examination is usually sufficient.

MRI of the primary tumour can be combined with an examination of the pelvic nodes. Tumour in virtually all cases involves superficial groin nodes first, and from there spreads to deep groin and pelvic nodes [22]. Certain features on MRI are highly specific for involvement—in particular, a fluid component indicating necrosis [6]—but are not common, and in general the most commonly used criterion to determine nodal involvement is short axis diameter. The fundamental limits to MRI are that (1) a small degree of infiltration will not significantly affect the size of a node, and (2) reactive nodal enlargement from local inflammation is particularly common in cancer of the penis, and the cause of nodal enlargement in 25–50% of palpable groin nodes at presentation [23, 24], although palpable nodes at follow-up are almost always malignant.

Table 1. Current and proposed classification for the local staging of penile cancer

<table>
<thead>
<tr>
<th>T stage</th>
<th>Current definition</th>
<th>Proposed definition*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tx</td>
<td>Cannot be assessed</td>
<td>Cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumour</td>
<td>No evidence of primary tumour</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
<td>Carcinoma in situ</td>
</tr>
<tr>
<td>T1</td>
<td>Non-invasive verrucous carcinoma</td>
<td>Non-invasive verrucous carcinoma</td>
</tr>
<tr>
<td>T1</td>
<td>Tumour invades subepithelial connective tissue</td>
<td>Tumour invades subepithelial connective tissue</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour invades corpus spongiosum or corpus cavernosum</td>
<td>Tumour invades corpus spongiosum</td>
</tr>
<tr>
<td>T3</td>
<td>Tumour invades urethra or prostate</td>
<td>Tumour invades corpus cavernosum</td>
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<tr>
<td>T4</td>
<td>Tumour invades other adjacent structures</td>
<td>Tumour invades adjacent structures (including prostate)</td>
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*According to Leijte et al [18].
The accuracy of size criteria is little better for pelvic nodes [26, 27].

**Priapism**

The first important distinction in priapism (prolonged, often painful erection) is the distinction between low- and high-flow states. Low flow is the commonest type and a form of compartment syndrome; as elsewhere in the body if untreated it leads to infarction and fibrosis [28]. It may be caused by several drugs, both therapeutic (including phosphodiesterase inhibitors and intracavernosal agents) and recreational, sickle cell disease, leukaemia and malignant infiltration, although in 30–50% of cases it is idiopathic [29]. In contrast, in high-flow priapism there is often an arteriolacunar fistula, the corpora are at least initially well oxygenated and there is usually no pain. The distinction between the two states is important, because low-flow priapism requires emergency treatment, but the diagnosis can normally be achieved either clinically, by measurement of blood gases in an aspirate, or on Doppler ultrasound [30].

There are two main indications for MRI. In low-flow priapism (Figure 5) the degree of corporal infarction may influence the decision to intervene, but more definitively if the corpora cavernosa are completely thrombosed after surgical attempts at a shunt, the best functional outcome may be achieved by early insertion of a penile prosthesis [31]. The degree of infarction on MRI correlates well with more established (but less anatomically accurate) methods of assessment such as Doppler and blood gas measurement, and with histology [31]. It is important to emphasise that perfusion of the corpora can be slow, and in addition to dynamically enhanced sequences to assess the patency of the cavernosal arteries, delayed sequences at 5 and 10 min after contrast are necessary to assess tissue viability.

In high-flow priapism, a fistula can be suspected on the dynamic post-contrast images when there is asymmetrical, early enhancement in one corpus, and is often seen as a focus of heterogeneous flow void on T2 weighted sequences (Figure 6). However, the combination of ultrasound [30] and angiography is more conventionally used, and is probably more sensitive than MRI, although no direct comparisons have been published.

**Penile fracture**

Penile fracture is a traumatic disruption of the tunica albuginea, often felt as a “snap” and usually associated with complete and rapid detumescence. It usually requires urgent surgery to prevent subsequent deformity and erectile dysfunction [32].

The penis should ideally be scanned in the anatomical position (to prevent confusing “kinking”) and without intracavernosal agents. The hallmark of a fracture is an interruption of the low-signal tunica albuginea (Figure 7), usually best seen on T2 weighted sequences. However, a T1 spin echo sequence may show the associated haematoma best, and in one small series was the only sequence that showed the fracture well in three patients; enhancement was not necessary [33].

Identification of the fracture may be useful to the surgeon because a localised exploration may then be performed, rather than an extensive subcoronal degloving procedure [34], which probably has a higher
post-operative morbidity [35]. MRI detects most fractures (seven out of nine in one series) [36], but should be used with caution when excluding the diagnosis, although there is little doubt that it is more sensitive than ultrasound and cavernosography [37]. Associated urethral injuries may be found in around a quarter of patients, and can be suspected on MRI [33], although urethrography remains the gold standard for their detection. MRI may also detect several pathologies that mimic penile fracture—in particular intracavernosal haematoma [34] and rupture of a superficial vein. Although we have been able to diagnose suspensory ligament rupture in one case [3], we do not know the accuracy of MRI in this condition.

Fibrosis and Peyronie’s disease

Fibrosis within the corpora or in the tunica albuginea may result from priapism, trauma or intracavernosal agents [36], but the commonest cause (occurring in around 3% of males) is Peyronie’s disease, defined clinically as a palpable plaque, usually with associated curvature [38], and possibly due to an aberrant healing response to repeated shear strains [39].

Tumescence with intracavernosal agents both improves the sensitivity for plaque and intracavernosal fibrosis (both seen as focal areas of low $T_2$ signal) and demonstrates associated angulation or waisting [40] (Figure 8). Although ultrasound may also demonstrate most plaques (and is superior for the detection of calcification) [41], it is probably a little less sensitive (with 67% and 61% of palpable plaques demonstrated on MR and ultrasound in one study [41]). Except at the base of the penis, MRI is still probably less sensitive for tunical plaque than clinical examination [41], and is probably not indicated routinely, although it can be useful for surgical planning.

One potential benefit of MRI is the demonstration of tunical enhancement, which in a small series [42] correlated with the initial inflammatory phase of Peyronie’s disease, when most surgeons advise against surgery. However, there is little correlation with pain—the most commonly used indicator of active disease [41]—and the significance of plaque enhancement remains uncertain.

Intracavernosal fibrosis (both in Peyronie’s disease and post traumatic) may be seen as areas of low-signal standing, but should not be overdiagnosed: some low-signal standing in the distal corpora cavernosa is seen in normal males [3].
Implants

Inflatable implants are all safe in 1.5 T systems but two malleable prostheses (OmniPhase and DuraPhase; Dacomed, Minneapolis, MN) containing metallic elements show fairly strong deflection in 1.5 T magnets [43] and should not be scanned.

Inflatable implants are well seen on T2 and STIR sequences [44], and we scan with the device inflated when possible. Although ultrasound is sufficient in many cases to check for adequate fluid in the device, several abnormalities are best seen on MRI. In particular, kinking from overlong cavernosal components can be hard to detect clinically, and in one study an anatomical abnormality was seen on MRI in all 14 patients with pain, apparent clinically in only 5 [45]. Crossover of cavernosal components is relatively rare, but well seen [3], as is aneurysmal dilation of the cavernosal components, which may lead to bulge and underinflation (Figure 9). The hypermobile glans, leading to a “supersonic transporter deformity” [46], can be diagnosed clinically or by ultrasound.

Infection occurs in around 2–4% of inflatable implants (although subclinical infection is more common) [47] and is often associated with periprosthetic fluid and stranding (seen in particular on STIR sequences) [45], but these findings should be interpreted with caution: some fluid does not necessarily imply infection, and stranding may be seen for several months after insertion. We have found that infections that are convincingly shown on MRI are usually clinically apparent as well.

Erectile dysfunction

Peyronie’s disease and penile fibrosis can both cause erectile dysfunction and can be imaged on MRI. However, clinical tests of nocturnal penile tumescence, response to phosphodiesterase inhibitors and intracavernosal agents, together with penile Doppler ultrasound remain the mainstay of imaging investigation [48, 49]. MR angiography may demonstrate the branches of the internal iliac vessels, and can be used to plan pelvic revascularisation, but is not of adequate resolution to show the penile vessels well, and conventional angiography is superior [50].

Urethrography

MR urethrography is technically feasible, but not necessarily straightforward. Compared with conventional urethrography it may sometimes show fistula, tumour, and spongiosis better [51], but similar benefits are seen with ultrasound urethrography of the anterior urethra [52] and both MRI and ultrasound are considerably more cumbersome than a fluoroscopic ascending and descending urethrogram, which is quick, demonstrates...
fistulas well and can show the posterior urethra during voiding.

MR urethrography can be performed with T2 sequences and saline or jelly [51] or T2 gradient echo sequences around an hour after injection of intravenous gadolinium [53]. One place where MRI may be useful is after pelvic trauma, where as well as delineating the urethral lumen it may demonstrate the anatomical abnormality and degree of prostatic displacement [54]; in particular, one report has suggested that it may be more accurate than conventional urethrography at determining the length of the obliterated segment [55], but it must be stressed that obtaining good urethral distension can be technically challenging. Periurethral enhancement correlates with inflammation but is as yet of uncertain clinical significance [56].

Miscellaneous conditions

MRI can be used to define the location of injected silicone [57] (in particular using silicone-suppressed sequences) or other substances, and T2 and STIR sequences will show dilated Cowper's ducts. Finally, MRI is an excellent tool for defining the anatomy in cases of ambiguous or abnormal genitalia [58].

Conclusion

MRI can contribute useful information for many different pathologies in the penis, but is in many cases not convincingly superior to clinical examination or ultrasound to justify its routine use. It is probably most useful in the investigation of the painful penile implant and acute low-flow priapism. It is sometimes useful for the local staging of penile cancer, for the localisation of penile fracture and for imaging complex cases of fibrosis.

References

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